

Search Results -

Term	Documents	
5.CLMUSPT,PGPB.	7	
(L5.CLM.).USPT,PGPB.	7	

	US Patents Full-Text Database US Pre-Grant Publication Full-Text Database		
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Database:	Derwent World Patents Index IBM Technical Disclosure Bulletins		7.
Search:	L6	A	Refine Search
	Recall Text Clear	_	I

Search History

DATE: Monday, September 22, 2003 Printable Copy Create Case

Set Name		Hit Count	
side by side			result set
DB=US	SPT,PGPB; PLUR=YES; OP=ADJ		
<u>L6</u>	L5.clm.	7	<u>L6</u>
<u>L5</u>	(vitaxin or alphav or alphavbeta3 or lm609)same (tumor\$ or tumour\$ or cancer\$)	146	<u>L5</u>
<u>L4</u>	L3.clm.	3	<u>L4</u>
<u>L3</u>	(vitaxin or alphav or alphavbeta3 or lm609)same (angiogenesis)	241	<u>L3</u>
<u>L2</u>	L1 and (vitaxin or alphav or alphavbeta3 or lm609)	5	<u>L2</u>
<u>L1</u>	brooks.in.	2944	<u>L1</u>

END OF SEARCH HISTORY

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Search Results - Record(s) 1 through 5 of 5 returned.

☐ 1. Document ID: US 20030176334 A1

L2: Entry 1 of 5

File: PGPB

Sep 18, 2003

PGPUB-DOCUMENT-NUMBER: 20030176334

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030176334 A1

TITLE: Methods and compositions useful for inhibition of angiogenesis

PUBLICATION-DATE: September 18, 2003

INVENTOR-INFORMATION:

NAME

CITY

STATE

COUNTRY

RULE-47

Brooks, Peter Cheresh, David A. Hollywood Encinitas CA CA US US

US-CL-CURRENT: 514/12; 435/226, 530/350

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMC Draw Desc Image

☐ 2. Document ID: US 20030113331 A1

L2: Entry 2 of 5

File: PGPB

Jun 19, 2003

PGPUB-DOCUMENT-NUMBER: 20030113331

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030113331 A1

TITLE: METHOD AND COMPOSITION FOR ANGIOGENESIS INHIBITION

PUBLICATION-DATE: June 19, 2003

INVENTOR-INFORMATION:

NAME

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US-CL-CURRENT: 424/155.1; 435/7.23, 530/388.25

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KMIC Draw Desc Image

☐ 3. Document ID: US 6500924 B1

L2: Entry 3 of 5

File: USPT

Dec 31, 2002

US-PAT-NO: 6500924

Record List Display

DOCUMENT-IDENTIFIER: US 6500924 B1

TITLE: Methods and compositions useful for inhibition of angiogenesis

DATE-ISSUED: December 31, 2002

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY

Brooks; Peter C. Hollywood CA Cheresh; David A. Encinitas CA Silletti; Steven A. San Diego CA

US-CL-CURRENT: 530/350; 435/975, 530/300, 530/382

Full Title Citation Front Review Classification Date Reference Sequences Attachments KMIC Draw Desc Image

☐ 4. Document ID: US 5766591 A

L2: Entry 4 of 5 File: USPT

Jun 16, 1998

US-PAT-NO: 5766591

DOCUMENT-IDENTIFIER: US 5766591 A

TITLE: Methods and compositions useful for inhibition of angiogenesis

DATE-ISSUED: June 16, 1998

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY

Brooks; Peter San Diego CA

Cheresh; David A. Cardiff CA

US-CL-CURRENT: 424/184.1; 424/185.1, 424/277.1, 530/300, 530/350, 530/380, 530/381,

<u>530/382</u>

Full Title Citation Front Review Classification Date Reference Sequences Attachments KMC Draw Desc Image

☐ 5. Document ID: US 5753230 A

L2: Entry 5 of 5 File: USPT May 19, 1998

US-PAT-NO: 5753230

DOCUMENT-IDENTIFIER: US 5753230 A

** See image for Certificate of Correction **

TITLE: Methods and compositions useful for inhibition of angiogenesis

DATE-ISSUED: May 19, 1998

INVENTOR - INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY

Brooks; Peter San Diego CA Cheresh; David A. Cardiff CA

US-CL-CURRENT: 424/158.1; 424/143.1, 424/145.1, 424/155.1, 424/174.1, 530/388.22,

530/388.24, 530/388.25, 530/388.8

Full Title Citation Front Review Classification Date Reference Sequences Attachments KMC Draw Desc Image **Generate Collection** Print **Term Documents** VITAXIN 234 **VITAXINS** 0 25 ALPHAV ALPHAVS 0 ALPHAVBETA3 37 0 ALPHAVBETA3S LM609 148 LM609S 0 (1 AND (ALPHAV OR ALPHAVBETA3 OR VITAXIN OR 5 LM609)).USPT,PGPB. (L1 AND (VITAXIN OR ALPHAV OR ALPHAVBETA3 OR 5 LM609)).USPT,PGPB.

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Previous Page Next Page

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L2: Entry 4 of 5

File: USPT

Jun 16, 1998

US-PAT-NO: 5766591

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TITLE: Methods and compositions useful for inhibition of angiogenesis

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INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY

Brooks; Peter San Diego CA Cheresh; David A. Cardiff CA

US-CL-CURRENT: 424/184.1; 424/185.1, 424/277.1, 530/300, 530/350, 530/380, 530/381,

530/382

CLAIMS:

What is claimed is:

- 1. A method of inducing solid tumor tissue regression in a patient comprising administering to said patient a composition comprising a therapeutically effective amount of an RGD-containing integrin .alpha..sub.v .beta..sub.3 antagonist sufficient to inhibit neovascularization of a solid tumor tissue.
- 2. The method of claim 1 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist preferentially inhibits fibrinogen binding to .alpha..sub.v .beta..sub.3 compared to fibrinogen binding to a.sub.IIb.beta..sub.3.
- 3. The method of claim 1 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is an RGD-containing polypeptide.
- 4. The method of claim 3 wherein said polypeptide is selected from the group consisting of c-(GrGDFV) (SEQ ID NO 4), c-(RGDFV) (SEQ ID NO 5), c-(RGDFv) (SEQ ID NO 7), and YTAECKPQVTRGDVF (SEQ ID NO 8), and a salt thereof.
- 5. The method of claim 4 wherein said salt is hydrochloride or trifluoroacetate.
- 6. The method of claim 1 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is a cyclic polypeptide.
- 7. The method of claim 1 wherein said administering comprises intravenous administration.
- 8. The method of claim 1 wherein said administering comprises transdermal administration.
- 9. The method of claim 1 wherein said administering comprises intramuscular administration.
- 10. The method of claim 1 wherein said administering comprises topical administration.
- 11. The method of claim 1 wherein said administering comprises subcutaneous administration.

- 12. The method of claim 1 wherein said administering comprises intracavity administration.
- 13. The method of claim 1 wherein said administering comprises peristaltic administration.
- 14. The method of claim 1 wherein said administering comprises one or more dose administrations daily for one or several days.
- 15. The method of claim 1 wherein said administering comprises a single dose intravenously.
- 16. The method of claim 1 wherein said amount is from about 0.1 mg/kg to about 300 mg/kg patient body weight.
- 17. The method of claim 1 wherein said administering is conducted in conjunction with chemotherapy.
- 18. The method of claim 1 wherein said solid tumor tissue is a carcinoma.
- 19. The method of claim 1 wherein said solid tumor tissue is a tumor of lung, pancreas, breast, colon, larynx or ovary.
- 20. The method of claim 1 wherein the patient is human.
- 21. A method of inhibiting solid tumor tissue growth undergoing neovascularization in a patient comprising administering to said patient a composition comprising a therapeutically effective amount of an RGD-containing integrin .alpha..sub.v .beta..sub.3 antagonist sufficient to inhibit solid tumor tissue growth.
- 22. The method of claim 21 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist preferentially inhibits fibrinogen binding to .alpha..sub.v .beta..sub.3 compared to fibrinogen binding to .alpha..sub.IIb .beta..sub.3.
- 23. The method of claim 21 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is an RGD-containing polypeptide.
- 24. The method of claim 23 wherein said polypeptide is selected from the group consisting of c-(GrGDFV) (SEQ ID NO 4), c-(RGDFV) (SEQ ID NO 5), c-(RGDFv) (SEQ ID NO 7), and YTAECKPQVTRGDVF (SEQ ID NO 8), and a salt thereof.
- 25. The method of claim 24 wherein said salt is hydrochloride or trifluoroacetate.
- 26. The method of claim 21 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is a cyclic polypeptide.
- 27. The method of claim 21 wherein said administering comprises intravenous administration.
- 28. The method of claim 21 wherein said administering comprises transdermal administration.
- 29. The method of claim 21 wherein said administering comprises intramuscular administration.
- 30. The method of claim 21 wherein said administering comprises topical administration.
- 31. The method of claim 21 wherein said administering comprises subcutaneous administration.
- 32. The method of claim 21 wherein said administering comprises intracavity administration.

- 33. The method of claim 21 wherein said administering comprises peristaltic administration.
- 34. The method of claim 21 wherein said administering comprises one or more dose administrations daily for one or several days.
- 35. The method of claim 21 wherein said administering comprises a single dose intravenously.
- 36. The method of claim 21 wherein said amount is from about 0.1 mg/kg to about 300 mg/kg body weight.
- 37. The method of claim 21 wherein said administering is conducted in conjunction with chemotherapy.
- 38. The method of claim 21 wherein said solid tumor tissue is a carcinoma.
- 39. The method of claim 21 wherein said solid tumor tissue is a tumor of lung, pancreas, breast, colon, larynx or ovary.
- 40. The method of claim 21 wherein the patient is human.
- 41. A method for treating a patient with inflammed tissue in which neovascularization is occurring comprising administering to said patient a composition comprising a therapeutically effective amount of an RGD-containing integrin .alpha..sub.v .beta..sub.3 antagonist.
- 42. The method of claim 41 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist preferentially inhibits fibrinogen binding to .alpha..sub.v .beta..sub.3 compared to fibrinogen binding to .alpha..sub.IIb .beta..sub.3.
- 43. The method of claim 41 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is an RGD-containing polypeptide.
- 44. The method of claim 43 wherein said polypeptide is selected from the group consisting of c-(GrGDFV) (SEQ ID NO 4), c-(RGDFV) (SEQ ID NO 5), c-(RGDFv) (SEQ ID NO 7), and YTAECXPQVTRGDVF (SEQ ID NO 8), and a salt thereof.
- 45. The method of claim 44 wherein said salt is hydrochloride or trifluoroacetate.
- 46. The method of claim 41 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is a cyclic polypeptide.
- 47. The method of claim 41 wherein said administering comprises intravenous administration.
- 48. The method of claim 41 wherein said administering comprises transdermal administration.
- 49. The method of claim 41 wherein said administering comprises intramuscular administration.
- 50. The method of claim 41 wherein said administering comprises topical administration.
- 51. The method of claim 41 wherein said administering comprises subcutaneous administration.
- 52. The method of claim 41 wherein said administering comprises intracavity administration.
- 53. The method of claim 41 wherein said administering comprises peristaltic administration.
- 54. The method of claim 41 wherein said administering comprises one or more dose

administrations daily for one or several days.

- 55. The method of claim 41 wherein said administering comprises a single dose intravenously.
- 56. The method of claim 41 wherein said amount is from about 0.1 mg/kg to about 300 mg/kg body weight.
- 57. The method of claim 41 wherein said inflammed tissue is arthritic.
- 58. The method of claim 41 wherein said arthritic tissue is present in a mammal with rheumatoid arthritis.
- 59. The method of claim 41 wherein said patient is human.
- 60. A method for treating a patient in which neovascularization is occurring in retinal tissue comprising administering to said patient a composition comprising a neovascularization-inhibiting amount of an RGD-containing integrin .alpha..sub.v .beta..sub.3 antagonist.
- 61. The method of claim 60 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist preferentially inhibits fibrinogen binding to a .alpha..sub.v .beta..sub.3 compared to fibrinogen binding to .alpha..sub.IIb .beta..sub.3.
- 62. The method of claim 60 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is an RGD-containing polypeptide.
- 63. The method of claim 62 wherein said polypeptide is selected from the group consisting of c-(GrGDFV) (SEQ ID NO 4), c-(RGDFV) (SEQ ID NO 5), c-(RGDFv) (SEQ ID NO 7), and YTAECKPQVTRGDVF (SEQ ID NO 8), and a salt thereof.
- 64. The method of claim 63 wherein said salt is hydrochloride or trifluoroacetate,.
- 65. The method of claim 60 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is a cyclic polypeptid.
- 66. The method of claim 60 wherein said administering comprises intravenous administration.
- 67. The method of claim 60 wherein said administering comprises transdermal administration.
- 68. The method of claim 60 wherein said administering comprises intramuscular administration.
- 69. The method of claim 60 wherein said administering comprises topical administration.
- 70. The method of claim 60 wherein said administering comprises subcutaneous administration.
- 71. The method of claim 60 wherein said administering comprises intracavity administration.
- 72. The method of claim 60 wherein said administering comprises peristaltic administration.
- 73. The method of claim 60 wherein said administering comprises one or more dose administrations daily for one or several days.
- 74. The method of claim 60 wherein said administering comprises a single dose intravenously.
- 75. The method of claim 60 wherein said amount is from about 0.1 mg/kg to about

- 300 mg/kg body weight.
- 76. The method of claim 60 wherein said retinal tissue is in said patient with diabetic retinopathy or macular degeneration and said, angiogenesis is retinal angiogenesis.
- 77. The method of claim 60 wherein said patient is human.
- 78. A method for treating a patient for restenosis in a tissue wherein smooth muscle cell migration occurs following angioplasty comprising administering to said patient a composition comprising a therapeutically effective amount of an RGD-containing integrin .alpha..sub.v .beta..sub.3 antagonist.
- 79. The method of claim 78 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist preferentially inhibits fibrinogen binding to .alpha..sub.v .beta..sub.3 compared to fibrinogen binding to .alpha..sub.IIb .beta..sub.3.
- 80. The method of claim 78 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is an RGD-containing polypeptide.
- 81. The method of claim 80 wherein said polypeptide is selected from the group consisting of c-(GrGDFV) (SEQ ID NO 4), c-(RGDFV) (SEQ ID NO 5), c-(RGDFv) (SEQ ID NO 7), and YTAECKPQVTRGDVF (SEQ ID NO 8), and a salt thereof.
- 82. The method of claim 81 wherein said salt is hydrochloride or trifluoroacetate.
- 83. The method of claim 78 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is a cyclic polypeptide.
- 84. The method of claim 78 wherein said administering comprises intravenous administration.
- 85. The method of claim 78 wherein said administering comprises transdermal administration.
- 86. The method of claim 78 wherein said administering comprises intramuscular administration.
- 87. The method of claim 78 wherein said administering comprises topical administration.
- 88. The method of claim 78 wherein said administering comprises subcutaneous administration.
- $89.\ \ \mbox{The method of claim }78$ wherein said administering comprises intracavity administration.
- 90. The method of claim 78 wherein said administering comprises peristaltic administration.
- 91. The method of claim 78 wherein said administering comprises one or more dose administrations daily for one or several.days.
- 92. The method of claim 78 wherein said administering comprises a single dose intravenously.
- 93. The method of claim 78 wherein said amount is from about 0.1 mg/kg to about 300 mg/kg body weight.
- 94. The method of claim 78 wherein said administering is conducted after agioplasty.
- 95. The method of claim 78 wherein said angioplasty is coronary angioplasty.
- 96. The method of claim 78 wherein said patient is human.

- 97. A method of reducing blood supply to a tissue required to support new growth of said tissue in a patient comprising administering to said patient a composition comprising a therapeutically effective amount of an RGD-containing integrin .alpha..sub.v .beta..sub.3 antagonist sufficient to reduce said blood supply to said tissue.
- 98. The method of claim 97 wherein said tissue is selected from the group consisting of a tumor tissue, an inflammed tissue, a tissue at risk for restenosis following angioplasty, and retinal tissue.
- 99. The method of claim 97 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist preferentially inhibits fibrinogen binding to .alpha..sub.v .beta..sub.3 compared to fibrinogen binding to .alpha..sub.IIb .beta..sub.3.
- 100. The method of claim 97 wherein said integrin .alpha..sub.v .beta..sub.3 antagonist is an RGD-containing polypeptide.
- 101. The method of claim 100 wherein said polypeptide is selected from the group consisting of c-(GrGDFV) (SEQ ID NO 4), c-(RGDFV) (SEQ ID NO 5), c-(RGDFV) (SEQ ID NO 7), and YTAECKPQVTRGDVF (SEQ ID NO 8), and a salt thereof.
- 102. The method of claim 101 wherein said salt is hydrochloride or trifluoroacetate.
- 103. The method of claim 97 said integrin .alpha..sub.v .beta..sub.3 antagonist is a cyclic polypeptide.
- 104. The method of claim 97 wherein said administering comprises intravenous administration.
- 105. The method of claim 97 wherein said administering comprises transdermal administration.
- 106. The method of claim 97 wherein said administering comprises intramuscular administration.
- 107. The method of claim 97 wherein said administering comprises topical administration.
- 108. The method of claim 97 wherein said administering comprises subcutaneous administration.
- 109. The method of claim 97 wherein said administering comprises intracavity administration.
- 110. The method of claim 97 wherein said administering comprises peristaltic administration.
- 111. The method of claim 97 wherein said administering comprises one or more dose administrations daily for one or several days.
- 112. The method of claim 97 wherein said administering comprises a single dose intravenously.
- 113. The method of claim 97 wherein said amount is from about 0.1 mg/kg to about 300 mg/kg body weight.
- 114. The method of claim 97 wherein said patient is human.

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End of Result Set

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L2: Entry 5 of 5

File: USPT

May 19, 1998

US-PAT-NO: 5753230

DOCUMENT-IDENTIFIER: US 5753230 A

** See image for Certificate of Correction **

TITLE: Methods and compositions useful for inhibition of angiogenesis

DATE-ISSUED: May 19, 1998

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY

Brooks; Peter San Diego CA Cheresh; David A. Cardiff CA

US-CL-CURRENT: <u>424/158.1</u>; <u>424/143.1</u>, <u>424/145.1</u>, <u>424/155.1</u>, <u>424/174.1</u>, <u>530/388.22</u>, <u>530/388.24</u>, <u>530/388.25</u>, <u>530/388.8</u>

CLAIMS:

What is claimed is:

- 1. A method for inhibiting angiogenesis in a solid tumor in a patient wherein cells of the tumor do not express levels of integrin .alpha..sub.v .beta..sub.3 detectable by immunohistochemistry comprising administering to said patient a composition comprising an angiogenesis-inhibiting amount of an anti-.alpha..sub.v .beta..sub.3 monoclonal antibody, whereby integrin .alpha..sub.v .beta..sub.3 expressed on the surface of vascular endothelial cells involved in said angiogenesis is contacted by said antibody resulting in inhibition in the blood supply to the tissue of said solid tumor.
- 2. The method of claim 1 wherein said anti-.alpha..sub.v .beta..sub.3 antibody inhibits binding of fibrinogen to integrin .alpha..sub.v .beta..sub.3 but does not substantially inhibit binding of fibrinogen to integrin .alpha..sub.IIb .beta..sub.3.
- 3. The method of claim 1 wherein said monoclonal antibody has the immunoreaction characteristics of monoclonal antibody LM609having ATCC accession number HB 9537.
- 4. The method of claim 1 wherein said angiogenesis-inhibiting amount is from about 0.1 mg/kg to about 300 mg/kg.
- 5. The method of claim 1 wherein said administering comprises intravenous administration.
- 6. The method of claim 1 wherein said administering comprises intrasynovial administration.
- 7. The method of claim 1 wherein said administering comprises transdermal administration.
- 8. The method of claim 1 wherein said administering comprises intramuscular administration.

- 9. The method of claim 1 wherein said administering comprises oral administration.
- 10. The methods of claim 1 wherein said administering is conducted in conjunction with chemotherapy.
- 11. The method of claim 1 wherein the patient is human.
- 12. The method of claim 11 wherein the antibody is humanized.
- 13. The method of claim 12 wherein the humanized antibody has the immunoreaction characteristics of monoclonal antibody LM609 having ATCC accession number HB 9537.
- 14. The method of claim 1 wherein the anti-.alpha..sub.v .beta..sub.3 antibody specifically binds integrin .alpha..sub.v .beta..sub.3 complex.
- 15. The method of claim 1 wherein the tumor is metastasized.
- 16. A method for inhibiting angiogenesis in a solid tumor in a patient wherein cells of the tumor do not express levels of integrin .alpha..sub.v .beta..sub.3 detectable by immunohistochemistry with monoclonal antibody LM609 having ATCC accession number HB9537, comprising administering to said patient a composition comprising an angiogenesis-inhibiting amount of anti-.alpha..sub.v .beta..sub.3 monoclonal antibody, whereby integrin .alpha..sub.v .beta..sub.3 expressed on the surface of vascular endothelial cells involved in said angiogenesis is contacted by said antibody resulting in inhibition in the blood supply to the tissue of said solid tumor.
- 17. The method of claim 16 wherein said anti-.alpha..sub.v .beta..sub.3 monoclonal antibody has the immunoreaction characteristics of monoclonal antibody <u>LM609</u> having ATCC accession number HB9537.
- 18. The method of claim 16 wherein said patient is human.
- 19. The method of claim 18 wherein said anti-.alpha..sub.v .beta..sub.3 monoclonal antibody is humanized.
- 20. The method of claim 19 wherein the humanized antibody has the immunoreaction characteristics of monoclonal antibody LM609 having ATCC accession number HB 9537.
- 21. The method of claim 16 wherein the anti-.alpha..sub.v .beta..sub.3 antibody specifically binds integrin .alpha..sub.v .beta..sub.3 complex.
- 22. The method of claim 16 wherein said administering comprises intravenous administration.
- 23. The method of claim 16 wherein said administering comprises intrasynovial administration.
- 24. The method of claim 16 wherein said administering comprises transdermal administration.
- 25. The method of claim 16 wherein said administering comprises intramuscular administration.
- 26. The method of claim 16 wherein said administering comprises oral administration.
- 27. A method for inhibiting angiogenesis in an inflamed, angiogenic tissue of a patient, comprising administering to said patient a composition comprising an angiogenesis-inhibiting amount of an anti-.alpha..sub.v .beta..sub.3 monoclonal antibody, whereby integrin .alpha..sub.v .beta..sub.3 expressed on the surface of vascular endothelial cells involved in said angiogenesis in said angiogenic tissue is contacted by said antibody resulting in inhibition in the blood supply to said angiogenic tissue.

- 28. The method of claim 27 wherein said monoclonal antibody has the immunoreaction characteristics of monoclonal antibody LM609 having ATCC accession number HB 9537.
- 29. The method of claim 27 wherein said patient is human.
- 30. The method of claim 29 wherein said antibody is humanized.
- 31. The method of claim 30 wherein the humanized antibody has the immunoreaction characteristics of monoclonal antibody $\underline{LM609}$ having ATCC accession number HB 9537.
- 32. The method of claim 27 wherein the anti-.alpha..sub.v .beta..sub.3 antibody specifically binds integrin .alpha..sub.v .beta..sub.3 complex.
- 33. The method of claim 27 wherein said tissue is arthritic.
- 34. The method of claim 33 wherein said arthritic tissue is present in a mammal with rheumatoid arthritis.
- 35. The method of claim 27 wherein said tissue is the retinal tissue of a patient with diabetic retinopathy and said angiogenesis is retinal angiogenesis.
- 36. The method of claim 27 wherein said administering comprises intravenous administration.
- 37. The method of claim 27 wherein said administering comprises intrasynovial administration.
- 38. The method of claim 27 wherein said administering comprises transdermal administration.
- 39. The method of claim 27 wherein said administering comprises intramuscular administration.
- 40. The method of claim 27 wherein said administering comprises oral administration.
- 41. The method of claim 27 wherein the tissue is retinal tissue the angiogenesis is retinal angiogenesis.